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Photoreduction of Triplet Thioxanthone Derivative by Azolium Tetraphenylborate: a Way to Photogenerate N-Heterocyclic Carbenes

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Although N-heterocyclic carbenes (NHCs) have brought profound changes in catalytic organic synthesis, their generation generally requires inert atmosphere and harsh conditions. To overcome these limitations, an air-stable NHC photogenerator has been developed involving two mild components: 1,3-bis(mesityl)imidazolium tetraphenylborate (IMeshBHPh4) and electronically excited isopropylthioxanthone (ITX). In this study, the photochemical mechanism is investigated through the accurate identification of the transient species and photoproducts. Electron transfer reaction between the excited triplet state of ITX and BPh4− is demonstrated as being the primary photochemical step. Nanosecond laser spectroscopy shows an efficient quenching and the formation of the expected ITX radical anion. The oxidized borane species is not observed, suggesting that this short-lived species could dissociate very rapidly to give phenyl radical – successfully identified using electron paramagnetic resonance – and triphenyloborane. As regards the final photoproducts, 1H and 13C NMR spectroscopy support the formation of the targeted NHC, 1,3-bis(2,4,6-trimethylphenyl)imidazol-2-ylidene (IMes), suggesting the occurrence of a subsequent proton transfer reaction between ITX radical anion and imidazolium cation (IMesh+). Gas chromatography-mass spectrometry reveals other products: biphenyl, isopropylthioxanthene and ITX. Their formation can be reconciled with a 2-step mechanism of photoinduced electron/proton transfer reactions. 11B NMR spectroscopy demonstrates that the main organoboron photoproduct is diphenyloborinic acid formed by oxidation of BPh4−. Due to its Lewis acidity, Ph2BOH can react with IMes to yield an NHC-boron adduct.

Introduction

Of high importance in photochemistry are aryloborates, which are four-coordinate boron compounds bearing a tetrahedral geometry, a formal negative charge, and at least one aryl substituent. Although their photochemical reactivity has been studied since the late 1970s,1,2 progress in this field has been made at the cost of intense debates,3 and many unknowns still remain.4 Photoactivity of aryloborates is organized around two major types of reactions (Figure 1): 1/ direct photolysis and 2/ photoinduced electron transfer (PET), in this latter case, the borate anion acts as one electron reducing agent. Recently, other reactions such as photochromic isomerization or photoelimination were described for aryloborate molecules with an N,C- or a C,C-chelate backbone.5

- Photoysis upon exposure to 254 nm irradiation (route 1) was focused on tetraarylborate compounds such as NaBPh4. The most accepted mechanism involves a di-π-borate rearrangement leading to a three-membered biradical anion, evolving towards a more stable borirane anion isomer. Most studies on direct irradiation attempted to identify photoproducts (biphenyl, 1-phenyl-cyclohexadiene mostly) and elucidate photochemical mechanism.1,6,7 Its utility in preparative photochemistry was demonstrated only recently.

Figure 1. Photochemical reactivity of aryloborate species. Route 1 is based on direct photolysis while Route 2 is a photoinduced electron transfer (PET) involving an electron acceptor (EA) sensitizer. The asterisk designates an excited state.
Observing that intermediate anion species could abstract acidic protons from water and alcohol, Sun et al. proposed in 2008 a tetraphenylborate salt containing as cation the conjugated acid of a nitrogen base (BH⁺) such as bicyclic guanidine TBD (TBDH)⁶. The base was photogenerated through photoinduced proton abstraction of BH⁺ during the photolysis at 254 nm. For example, TBDH’BPh₄⁺ was used as photobase generator (PBG) for the organocatalyzed ring-opening polymerization of cyclic esters⁶ or thiol-epoxy polymerization.⁹ Several authors added an aryl ketone derivative to extend absorption to near UV-Vis range via a supposed triplet-triplet energy transfer.¹⁰–¹³

Due to their relatively low oxidation potential,¹⁴ arylborates such as Ph₃B⁺ or triarylalkylborate salts (Ph₃BR⁺) were also used as electron donor in inter- and intra-PET reactions (route 2) with cationic or neutral electron acceptor (EA) sensitizers such as carbocyanine,¹⁵ fullerene,¹⁶ coumarine,¹⁷ fluorone,¹⁸ benzophenone derivatives.¹⁹ A boranyl radical is formed, which in turn undergoes rapid cleavage yielding a triphenylborane and a phenyl (or alkyl) radical. Only alkyl radicals have been applied for initiation of radical polymerizations,¹⁷ or alkylation of the EA.²⁰

Recently, we proposed another application for photoinitiated oxidation of arylborates, the photogeneration of N-heterocyclic carbene (NHC) from an azolium arylborate salt (NHCH⁺BPh₄⁺) (Figure 2).²¹ When paired with an EA sensitizer such as isopropylthioxanthone (ITX), an electron transfer from the borate (Ph₃B⁺) to the excited triplet state of ITX (‘ITX³’) may occur as described above. Unlike the previous case, the azolium cation (NHCH⁺) enables a subsequent step of proton abstraction with ITX radical anion ITX⁻ (1) to take place, yielding the expected free NHC (2).²¹ Our recent investigations proved that it was possible to form iMes and its saturated analogue SIMes, the two most employed NHCS.²¹ The utility of this two-component NHC photogenerator NHCH⁺BPh₄⁺/ITX was also demonstrated in photopolymerization reactions to form polyurethane, polyester and polynorbornene.²₂ Indeed, although NHCS have brought profound changes in catalytic organic synthesis,²³–²⁵ they generally require inert atmosphere and harsh conditions for their generation. As a result, an air-stable system based on mild components, able to generate NHC on demand and on simple UV exposure has the potential to significantly simplify implementation of the broad spectrum of NHC-catalyzed reactions. While it is true that thermally latent NHC precursors also exist,²⁶,²⁷ photochemically produced NHCS have net advantages including ambient temperature reaction, thermal stability and the possibility to finely tune NHC concentration upon adjusting the energetic dosage.

Though the mechanistic hypothesis for photoreduction of thioxanthone derivative by arylborates is basically acceptable, conclusive evidences in support of a coupled electron transfer/proton abstraction are required. With exception of the photogenerated NHC molecule (2), little is known about the transient species – ITX⁻, ITX⁻ (1), ITXH⁺ (3), Ph₃B⁺ (4), Ph⁺ (5) – which are supposed to form and the other putative products such as Ph₃B (6) and the various thioxanthone derivatives. To provide conclusive evidences in support of our mechanism, it is important to clearly establish the identity of transient species and photoproducts. The structure of the organoboron compound(s) formed is particularly important given BPh₃ is likely to form ate-complexes with NHCS, with strong implications for carbene reactivity.²⁶ Indeed, trivalent boron species typically have Lewis acid characteristics, while NHCS behave as Lewis base. Motivated by this situation, this paper investigates the identity of transient species and photoproducts generated by the irradiation of 1,3-bis(mesityl)imidazolium tetraphenylborate (iMesH⁺BPh₄⁺, Figure 2) with ITX to form the NHC 1,3-bis(mesityl)imidazol-2-ylidine (iMes). To this purpose, a range of techniques was used: nanosecond laser spectroscopy and electron paramagnetic resonance (EPR) for the detection of the transient species, as well as gas chromatography-mass spectrometry (GC-MS) and ¹¹B NMR spectroscopy for the identification of the photoproducts.

![Figure 2](image_url)

**Figure 2.** Photochemical pathway for the generation of NHC from a mixture of NHCH⁺BPh₄⁺ and ITX.
Experimental

Materials

1,3-Bis(2,4,6-trimethylphenyl)imidazolium chloride (IMesHCl, 98.0 %, Aldrich), sodium tetrathyenylborate (NaBPh₄, 99.5 %, Aldrich), N-tet-butyl-α-phenyltrinitrene (PBN, 98.0 %, Aldrich), diphenyl(2,4,6-trimethylbenzoyl)phosphine oxide (TPO, 98 %, Aldrich), 2-isopropylthioxanthone (ITX, analytical standard, Aldrich), 1,3-bis(2,4,6-trimethylphenyl)imidazol-2-ylidene (IMes, 97%, Aldrich) and carbon disulfide (CS₂, anhydrous, Aldrich) were used as received unless otherwise mentioned. Acetonitrile-d₃ (ACN-d₃, 99.8 % D, Aldrich), tetrahydrofuran-d₈ (THF-d₈, 99.5 % D, Eurisotop), acetonitrile (ACN, HPLC grade, VWR) and ethanol (EtOH, HPLC grade, VWR) were dried over 4 Å molecular sieves before use.

Synthesis of IMesH²⁺BPh₄⁻

The IMesH²⁺BPh₄⁻ was synthesized as described previously.²¹ ¹H NMR (300 MHz, THF-d₈), IMesH²⁺BPh₄⁻: δ ppm: 2.03 (s, 12H, 4 × α-CH₂), 2.37 (s, 6H, 2 × p-CH₃), 6.62 – 6.66 (t, 4H, J = 6 Hz, 4 × ArH), 6.75 – 6.80 (t, 8H, J = 7.5 Hz, 8 × ArH), 7.13 (s, 4H, 4 × ArH), 7.15 (s, 2H, 2 × NCH) 7.22 – 7.26 (br, 8H, 2 × 8 × ArH), and 8.70 (s, 1H, NCHN).

Characterization methods

Nanosecond laser spectroscopy. All nanosecond time-resolved transient absorption spectra were performed in a 1 cm optical path length and monitored by Edinburgh Instruments LP920 laser flash photolysis spectrometer. The solutions were excited at 90° from the probe beam by a Q-switched nanosecond Nd/YAG laser (λexc = 355 nm, 8 ns pulse duration; energy reduced down to 5 mJ per pulse from Continuum (Surelite II–10)). A filter was used to remove the residual excitation light at 355 nm (Schott GG385). As preliminary step, all acetonitrile solutions were purged with nitrogen for 5 min prior to measurement. The decay of the triplet-triplet state absorption of ITX (1 × 10⁻³ M) at 600 nm was measured and follows a first-order kinetics. The temporal change of triplet concentration [ITX⁺] can be represented in equation (1):

\[ [\text{ITX}^+] (t) = [\text{ITX}^+] (0) \times e^{-t/\tau_0} \]  

where \( \tau_0 \) is the lifetime of the ITX triplet without quencher.

Abs. (t, 600 nm) = Abs. (0, 600 nm) × e⁻ᵗ/τ₀  

Exponential fitting of the experimental decay Abs.(t, 600 nm) provides \( \tau_0 \) (Equation (2)). In presence of a triplet quencher (IMesH²⁺BPh₄⁻, NaBPh₄ or IMesHCl), pseudo-first-order kinetic equation can be applied to extract a triplet lifetime \( \tau \) that depends on quencher concentration \([Q]\). Note that all the triplet lifetimes were determined by using a double exponential equation in the time range 0 – 40 µs the longer time-constant being due to the formation of the ITX radical anion at the monitored wavelength. By measuring \( \tau \) at different \([Q]\), the quenching rate constants, \( k_q \) can be obtained by the linear plot based on the Stern-Volmer equation (3).²⁹

\[ \tau^{-1} = \tau_0^{-1} + k_q [Q] \]  

Electron paramagnetic resonance (EPR). EPR measurements were performed with a Bruker Elexsys E500 spectrometer with X band frequency in continuous wave (around 9.8 GHz) at room temperature. Spectra were recorded with a modulation amplitude of 1 G, a modulation frequency of 100 kHz and a microwave power of ~ 2 mW. Both Bruker WIN-EPR and SimFonia software were used to note the spectra and carry out the simulation afterwards. In a typical experiment, an acetonitrile solution (1 mL) containing IMesH²⁺BPh₄⁻ (9.18 mg, 5 equiv.), ITX (1.27 mg, 3 equiv.) and PBN (0.54 mg, 1 equiv.) was degassed prior to transfer into an aqueous EPR cell. The cell was irradiated for different times (0 s, 30 s and 60 s) with a 365 nm LED light-guide (LC-L1V3, Hamamatsu, 65 mW·cm⁻²) then an EPR spectrum was acquired.

¹H NMR and ¹³C NMR spectroscopy. All ¹H NMR and ¹³C NMR spectra were recorded in appropriate deuterated solvents with tetramethylsilane (TMS) as the internal reference on a Varian Mercur 300 MHz. In a typical measurement, IMesH²⁺BPh₄⁻ (9.18 mg, 0.015 mmol, 3 equiv.) and ITX (1.27 mg, 0.005 mmol, 1 equiv.) were charged into a borosilicate NMR tube. The mixture was dissolved in 0.3 mL of acetonitrile-d₈ (0.03 M, relative to IMesH²⁺BPh₄⁻). Before exposure to a 365 nm LED light spot during 5 min (see details in previous section), the NMR tube was capped with a rubber septum and degassed with N₂. The as-irradiated tube was then analyzed by ¹H NMR. Excess amount of CS₂ (0.02 mL) was added subsequently into the tube. It caused a sudden color change from yellow to red followed by a gradual precipitation. The suspension media was kept for 24 h, and the solid was collected by vacuum filtration and air dried for 24 h. ¹H NMR and ¹³C NMR measurements were performed in DMSO-d₆ to confirm the formation of the IMes-CS₂ zwitterion adduct.

¹²B NMR spectroscopy. All ¹¹B NMR were recorded in THF-d₈ with BF₃.Et₂O as the internal reference on a Varian Bruker Avance NEO 500 MHz. All solutions were prepared under argon using a glove box technique. Prior to utilization, quartz NMR tubes were dried under vacuum overnight at 80°C. The THF-d₈ solvent was dried over molecular sieves.
Gas chromatography-mass spectrometry (GC-MS). The GC instrument is a Shimadzu GC-2010 model completed with the QP-2010 mass spectrometer system. Helium is the gas vector, used at the constant linear velocity of 25 cm s⁻¹. The sample was injected into a Supelco® BP-X5 column (0.15 mm of diameter, 25 m of length and 0.25 µm of film thickness) by using split mode with a ratio of 75:25. The temperature of the injector is maintained at 330°C. The column temperature was heated from 70°C to 340°C in two subsequent steps: the first heating state to 250°C with heating rate of 15°C min⁻¹ and the second to 340°C at the rate of 4°C min⁻¹. Then, the sample was ionized using an electronic ionization source heated at 200°C. The mass spectrum was recorded from 4 min to end of the program with scan mode from 50 m/z to 600 m/z. Mass Spectra of each peak were compared with spectra of NIST05 and NIST05s databases to identify compounds. In a typical experiment, 2 mL of acetonitrile solution including IMesH⁺BPh₄⁻ (1.71 mg, 0.003 mmol, 3 equiv.) and ITX (0.24 mg, 0.001 mmol, 1 equiv.) were placed into a UV quartz cuvette (1 cm optical path) and closed by a rubber septum. The cuvette was degassed with N₂ during 5 min then immediately exposed to a 365 nm LED spot light (65 mW·cm⁻²) for 5 min while keeping the stirring. Finally, the irradiated solution was transfer into Agilent 2 mL glass vial with a screw top (PTFE septum) under inert condition for GC-MS analysis.

Results and discussion

Assignment of transient species

Nanosecond laser spectroscopy

To identify the intermediate species formed in the 2-component system ITX/IMesH⁺BPh₄⁻ (Figure 2), excitation transfer reactions of ITX with different quenchers were recorded by means of a nanosecond laser spectroscopy. Laser flash exposure was performed at 355 nm in deaerated and anhydrous acetonitrile, where only the sensitizer (ITX) absorbs the light. Given the short lifetime of ITX singlet state as well as its high intersystem crossing quantum yield, ITX is assumed to interact mainly via its triplet state. As can be seen in Figure 3a, the transient absorption spectra of ITX alone shows an absorption maximum at 630 nm, fully consistent with the ITX triplet (³ITX*) in acetonitrile reported in the literature.30 Slow decrease of the transient absorption was attributed to the deactivation process of the ³ITX* with a lifetime of 5.7 µs at an initial ITX concentration of 10⁻⁵ M. To study the effect of the azolium cation, the reactivity of the triplet state was evaluated in presence of IMesH⁻Cl⁻ since no quenching is expected from Cl⁻ anion.30 The absorption at 630 nm was weakly quenched when this imidazolium salt was added to a deoxygenated solution of ITX. A low rate constant (4.7 × 10⁸ M⁻¹ s⁻¹) was determined by constructing a Stern-Volmer plot of the triplet lifetime as a function of [IMesH⁻Cl⁻], with the triplet being monitored at 600 nm (see experimental section for details). A typical decay experimental trace for the triplet decay of ITX in presence of IMesH⁻Cl⁻ is given in Figure 4 (trace a), showing clean first-order kinetics. The existence of very weak interactions is consistent with the fact that photooxidation of ³ITX* by IMesH⁻ is thermodynamically unfavorable (ΔG° = + 0.74 eV)32 and that the triplet energy transfer between ITX (ET = 2.77 eV) and IMesH⁻ (ET = 3.34 eV) exhibits a fairly endothermic character (Fig. S1 in Supporting Information). It can thus be concluded that the azolium cation plays a minor role in the primary photochemical reaction.

Similarly, triplet quenching experiment was carried out with NaBPh₄ to investigate the specific role of the tetraarylborate anion given that Na⁺ cannot quench or donate proton. As can be seen in Figure 4 (trace b), the decay rate was much faster, illustrating the more efficient quenching of ³ITX* by BPh₄⁻. Additionally, it can be noticed that the decay trace does not return to zero in this case; deviation to first order kinetics was attributed to the fact that the triplet absorption probably overlaps with another species. To clarify this point, Figure 3b shows the temporal evolution of the transient absorption spectrum of ITX/NaBPh₄. After 100 ns, it seems that the narrow 630 nm peak due to triplet absorption has disappeared and has been replaced by a new broad absorption in the region of 560 – 680 nm. Subsequent spectra show the slow decay of this broad peak. We hypothesized that this transient species could be the radical anion ITX⁻ generated after photoreduction of ³ITX* by BPh₄⁻. To support our assignment, we rely first on the fact that the electron transfer for this reaction is slightly exothermic (ΔG° ≅ − 0.13
ev), because of the high oxidation potential of the donor anion BPh$_4^-$. Second, triplet energy transfer, which is the second possible interactions process between ITX ($E_T = 2.77$ eV) and BPh$_4^-$ ($E_T = 3.64$ eV) (Fig. S1), is not energetically favorable. Third, our transition absorption spectrum strongly resembles that of thioxanthone radical anion observed by Schuster et al., who reported also a similar broad absorption at 650 nm. This lends additional confidence to the involvement of $\tilde{3}$TX$^*$ and the operation of a charge transfer mechanism. However, there is no additional band in the transient spectrum that might be attributed to the boranyl radical 4 or other boron species derived from the oxidation of tetraphenyl borate anion. In borate salts, it is well established that back electron is not significant but that boronyl radical rearranges during the laser pulse. Therefore, the failure to observe Ph$_3$B$^-$ (4) indicates that a very short-lived character as this radical probably dissociates or rearranges during the laser pulse.

Having investigated the individual role of IMesH$^+$ and BPh$_4^-$, quenching experiment was then carried out using IMesH$^+$BPh$_4^-$ (Fig. 4, trace d). The similarity of quenching rate obtained by this quencher ($4.9 \times 10^{-7}$ M$^{-1}$ s$^{-1}$) and NaBPh$_4$ ($4.1 \times 10^{-7}$ M$^{-1}$ s$^{-1}$) is well indicative that the photoreduction of ITX by BPh$_4^-$ is the primary photochemical process. Additionally, a broad feature characteristic of ITX$^+$ was also observed in the region of 560 – 680 nm (Fig. S2). However, there is no additional absorption band in the 380 – 430 nm region that could be assigned to a thioxanthone ketyl radical 3 (ITX$^+$) derived from subsequent proton transfer step of ITX$^+$ with IMesH$^+$. This result can be understood on the basis of a much slower proton transfer step compared to electron transfer. The only indication for protonation of the radical anion 1 is the shorter lifetime of ITX$^+$ in presence of IMesH$^+$BPh$_4^-$ (29.5 ms) compared to NaBPh$_4$ (40.1 ms). A final quenching experiment was also carried out using IMes, the targeted NHC. A significant acceleration of the decay at 600 nm is observed, leading to an increased quenching rate ($7.2 \times 10^8$ M$^{-1}$ s$^{-1}$). Such a result is consistent with the stronger donor properties of IMes ($E_{ox} = -0.84$ eV) compared to BPh$_4^-$ ($E_{ox} = 0.94$ eV). It also highlights a more complex mechanism where photogenerated NHC can also interact with triplet ITX and compete with BPh$_4^-$ for charge transfer.

**Figure 4.** Decays of $\tilde{1}$TX$^*$ at 600 nm in the presence and absence of: (a) IMesH$^+$Cl$^-$, (b) NaBPh$_4^-$, (c) NaBPh$_4^-$ (under O$_2$) and (d) IMesH$^+$BPh$_4^-$. (ITX) $= 10^{-4}$ M, [IMesH$^+$Cl$^-$] = [NaBPh$_4^-$] = 6 $\times$ 10$^{-4}$ M, respectively, recorded in N$_2$-saturated acetonitrile.

**Figure 5.** Electron paramagnetic resonance (EPR) spectra of radical adduct from a solution: (a) ITX – NaBPh$_4^-$ – PBN and (b) ITX – IMesH$^+$BPh$_4^-$ – PBN after 60 s of irradiation. (ITX) $= 5 \times 10^{-4}$ M, [IMesH$^+$BPh$_4^-$] $= 1.5 \times 10^{-4}$ M, [NaBPh$_4^-$] $= 1.5 \times 10^{-4}$ M and [PBN] $= 3 \times 10^{-5}$ M in acetonitrile.
Assignment of photoproducts

Using $^1$H NMR spectroscopy, we found that irradiation of an oxygen-free acetonitrile-$d_3$ solution containing IMesH$^+$BPh$_3$ (1 equiv.) and ITX (3 equiv.) at 365 nm resulted in a complete deprotonation at C2 position ($H_e$, $\delta$ = 8.70 ppm) of the imidazolium cation (Figure 6). This result clearly suggests the generation of IMes (2). Further evidence was given upon adding CS$_2$ to the as-irradiated medium. A red precipitate forms immediately that was straightforwardly assigned to IMes-CS$_2$ using $^1$H and $^{13}$C NMR analysis (Fig. 5S). With these results and the identification of ITX$^-$ described above, it is thus possible to formulate a tentative second step where NHC are created through proton abstraction of IMes$^+$ cation by ITX$^-$. Two methods to determine NHC concentration that were described in previous publications (acid/base titration) and $^1$H NMR analysis of NHC-carbodiimide adduct) revealed a similar IMes$^+$ conversion of 50% after 5 min irradiation, which is in conflict with NMR data. A partial conversion means that IMes coexists in the reaction medium with its conjugated acid IMes$^-$ and may form H-bonded bis(carbene)-proton adduct. Formation of this C···H–C type adduct has been already observed spectroscopically, e.g. UV, NMR. In our case, evidence for the formation of bis(carbene)-proton adduct is reflected by the slight shift to lower resonance of methylene protons $H_e$ ($\delta$ = 2.07 ppm) and $H_e$ ($\delta$ = 2.37 ppm) observed after irradiation (trace b) compared to the initial state (trace a). Similar upfield shift was evidenced when adding 1 equiv. of free IMes to 9 equiv. of IMesH$^+$BPh$_3$ (Fig. S6). In this spectrum, the bridging proton $H_e$ ($\delta$ = 8.70 ppm) of the C···H–C complex disappeared entirely due to H-bonding. This confirms that protonation degree cannot be used to quantitatively assess the NHC yield.

GC-MS

To move forward in the identification of other photoproducts, the freshly irradiated photolysis medium was analyzed by GC-MS. Three products were detected: biphenyl, isopropylthioxanthene and ITX (Fig. S7 and Table S1). Biphenyl was obtained in good yield and its formation was also observed in other (photo)oxidation experiments of BPh$_3$ anion by iron (II) complex, 1,4-dicyanophthalene or coumarin derivative. However, the formation of biphenyl upon coupling of phenyl radical (detected by EPR) is unlikely. Labelling studies revealed indeed that two phenyl groups of the biphenyl come from the same borate, suggesting that the biphenyl was formed from carbons that were bound to boron. It is therefore reasonable to assume that biphenyl comes from triphenylborane. The most plausible explanation is that Ph$^+$ reacts rapidly with Ph$_3$B to produce Ph–Ph and Ph$_3$B$^-$.

In contrast, isopropylthioxanthene and ITX result from the ensuing ground-state chemistry and might derive from ITXH$^+$ (3), the thioxanthyl ketyl radical that is formed after the protonation process of ITX$^-$, and that we were not able to detect by nanosecond spectroscopy. As described in Figure 7, the ketyl radical 3 may undergo two types of reaction: first, a disproportionation, giving ITX and isopropylthioxanthol; second, a dimerization leading to dithioxanthyl pinacol derivative. However, except ITX, none of these thioxanthene reduction products were detected by GC-MS. This is apparently due to their thermal instability in the GC injection port as established by Schuster et al. Isopropylthioxanthol is known to disproportionate upon heating to form ITX and isopropylthioxanthene. Additionally, pinacol can be easily oxidized under air to regenerate ITX and isopropylthioxanthene. Additionally, pinacol can be easily oxidized under air to regenerate ITX and isopropylthioxanthene. Note also that 1-phenyl-1,4-cyclohexadiene is not detected while it is a major product of the direct photolysis of BPh$_3$ in presence of H donors. Its absence confirms that energy transfer from ITX to BPh$_3$ is not the predominant mechanism.

Figure 7. Side reactions involving ITX ketyl radical.

Figure 6. $^1$H NMR spectra change of ITX (0.07 M) and IMesH$^+$BPh$_3$ (0.21 M) in ACN-$d_3$ prior to UV exposure (a) and after 5 min irradiation (b).
In arylboron photochemistry, the identification of boron photoproducts represents a major challenge. Boron products can be involved in complex reactions where both the nature and the role of the boron species can be hard to determine. Additionally, their proportion generally differ depending on whether irradiation was performed under N₂ or air. As emphasized in Figure 2, the oxidized borate is assumed to evolve in triphenylborane, with possibly subsequent degradation reactions. In our case, the photogenerated Ph₃B could for example form ate-complexes with the photogenerated NHC Lewis base. ¹¹B NMR spectroscopy remains the best tool to identify the boron species. In the case of photoinitiated oxidation of arylborates, the NMR identification has rarely been studied, while it can serve to provide insights into the mechanism. Nevertheless, precise identification of the groups attached to boron is not trivial owing to the limited structural information that can be derived from ¹¹B NMR data.

Figure 8 shows the ¹¹B NMR spectrum of an argon saturated solution of ITX/IMesH⁺BP₃⁻ (1/1 equiv.) prior (a) and after 10 min irradiation (b). As expected, spectrum a exhibits only a narrow singlet attributed to BP₃⁻ anion (δ = -6.3 ppm). This chemical shift is characteristic of tetraorganoborate anions experiencing a strong deshielding. In the spectrum b obtained after irradiation, the residual resonance of BP₃⁻ is visible as well as two new features: an intense peak at 1.1 ppm, and a broad and much weaker signal at 5.9 ppm. Acting on the assumption that these species might derive from triphenylborane (6), a set of experiments using BP₃ was implemented. The ¹¹B NMR spectrum of Ph₃B (shown in Fig. S8) reveals a single and characteristic singlet at 50.3 ppm that is absent in the photolysis medium. In addition, the irradiation of Ph₃B with or without ITX (1 equiv.) did not cause any change in the spectrum. In contrast, the addition of IMes (1 equiv.) resulted in the formation of a white precipitate and a downfield chemical shift (-9 ppm, see Fig. S9) characteristic of NHC-borane species, that was not observed in spectrum b. Though the above experiments do not necessarily demonstrate that Ph₃B has not been formed, it is interesting to note that both free Ph₃B or Ph₃B-IMes adducts are not present in the photolysis medium, and that degradation reaction with triplet excited ITX cannot be advanced as explanation. As hypothesized in the last section, it is thus more reasonable to conclude that Ph₃B may react with Ph⁺ to form Ph₃B⁺.

To shed further light into the identity of these boron-containing species, CS₂ was added to the photolysate solution (ITX/IMesH⁺BP₃⁻) just after irradiation (Figure 8, spectrum c). This led to immediate conversion of the intermediate at 1.1 ppm to a new broad resonance at 45.2 ppm while the position of the second resonance at 5.9 ppm remained unchanged. The change of chemical shift suggests that a NHC-borane species was initially present. The addition of CS₂ caused its

**Figure 8.** ¹¹B-NMR spectra of a mixture of ITX (0.03 M) and IMesH⁺BP₃⁻ (0.03 M) in THF-d₈: (a) prior irradiation, (b) after 10 min UV exposure, and (c) after adding CS₂ into medium. Irradiation conditions: LED 365 nm, 65 mW·cm⁻².

**Figure 9.** ¹¹B-NMR spectra in non-dried THF-d₈: (a) Hydrolyzed Ph₃B, (b) Addition of IMes into hydrolyzed Ph₃B. The asterisk is used to designate an impurity of IMes.

**Figure 10.** Mechanism accounting for the formation of diphenylborinic acid (Ph₂BOH).
dissociation and the subsequent formation of the more stable IMes-CS₂ zwitterionic adduct. By this means, a free boron species was released whose value of $^{11}$B chemical shift at 45.2 ppm can be useful for its assignment. Observed $^{11}$B NMR spectra are consistent with the exact chemical shift of diphenylborinic acid Ph₂BOH. Consequently, the singlet signal at 1.1 ppm could be ascribed to IMes-Ph₂BOH adduct present in the photolysis solution. However, the second weak signal at 5.9 ppm could not be assigned. Notably, this second boron-containing complex is not able to form ate-complex with IMes due to the absence of change in the chemical resonance when CS₂ is introduced. To provide further evidence for this assignment, Ph₂BOH was generated in situ by moderate hydrolysis of the Ph₂B in non-dried THF-$d_8$ upon exposing the solution in the NMR tube to atmosphere. This solution revealed two broad signals centered at 44.9 ppm (Ph₂BOH) and 50.6 ppm (residual Ph₂B) in the $^{11}$B NMR spectrum (Figure 9). Addition of IMes into the medium caused the two signals to shift at ~9 ppm (Ph₂B-IMes adduct) and 1.1 ppm (IMes-Ph₂BOH adduct). This latter is visible in the spectrum b of Figure 8, lending further confidence in our assignment. The last issue to be settled concerns the mechanism leading to the formation of Ph₂BOH. As described in Figure 10, we postulate that the formation of Ph₂BOH may relate to the autoxidation of Ph₂B species. Reaction of Ph₂B with Ph⁺ yields Ph₂B⁺ and Ph₂. Ph₂B⁺ can react subsequently with traces of oxygen (even if argon purged NMR tube were used) to form the peroxyl radical Ph₂BOO⁺. New reaction with Ph₂B might form the peroxide species Ph₂BOOB₃B₂, easily decomposed in Ph₂BO⁺. Ph₂BOH could be formed by hydrogen abstraction with proton donor or after hydrolysis with water.

Conclusions

The photochemical mechanism underlying the release of NHC upon irradiation of IMesh⁺BPh₄⁻ with ITX has been clarified. As summarized in Figure 11, the first step is an electron transfer ($k_{d} \approx 4.9 \times 10^{5} \text{ M}^{-1} \text{ s}^{-1}$) between electronically excited ITX and borate anion BPh₄⁻ acting as electron donor to yield the ion pair IMesh⁺/ITX⁻ and the boranyl radical Ph₃B⁺. This latter is subjected to a rapid decomposition into Ph⁺ and Ph₃B. A subsequent reaction of the ion pair is proton abstraction of IMesh⁺ by ITX⁻ to form the IMes (NHC) and the ketyl radical (ITXH⁺). Of high importance for NHC reactivity was the fate of organoboron products. Ph₂B and Ph⁺ were proved to undergo a multi-step oxidation reaction, yielding biphenyl and diphenylborinic acid. Ph₂BOH was found to react with the photogenerated IMes to give an IMes-Ph₂BOH ate-complex. While it is difficult to avoid the formation of NHC-boron adduct, our recent investigation showed that such complex did not prevent the activity of its NHC photogenerator in ring-opening methathesis polymerization or for synthesis of polyurethane, but could have a detrimental effect on other reactions. Knowledge about the precise photochemical mechanism offers now the possibility for improving the photoinitiating system in a rational way, and create a new generation of NHC photogenerator.

Conflicts of interest

There are no conflicts to declare.

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Figure 11. General mechanistic pathway involved in the formation of IMes during the photolysis of a mixture ITX/IMesh⁺BPh₄⁻.